

High quality reference genome of drumstick tree (*Moringa oleifera* Lam.), a potential perennial crop

TIAN Yang^{1,10,13†}, ZENG Yan^{4†}, ZHANG Jing^{8†}, YANG ChengGuang⁹, YAN Liang^{1,5},
WANG XuanJun¹³, SHI ChongYing², XIE Jing³, DAI TianYi², PENG Lei², ZENG HUAN Yu¹,
XU AnNi¹, HUANG YeWei¹³, ZHANG JiaJin^{11,12}, MA Xiao¹³, DONG Yang^{7,10},
HAO ShuMei^{6*} & SHENG Jun^{13*}

¹College of Life Sciences, Jilin University, Changchun 130012, China;

²College of Food Sciences, Yunnan Agricultural University, Kunming 650201, China;

³College of Animal Sciences, Yunnan Agricultural University, Kunming 650201, China;

⁴College of Life Sciences, University of Chinese Academy of Sciences, Beijing 100049, China;

⁵Pu'er Institute of Pu-er Tea, Pu'er 665000, China;

⁶School of Agriculture, Yunnan University, Kunming 650091, China;

⁷Faculty of Life Science and Technology, Kunming University of Science and Technology, Kunming 650093, China;

⁸College of Life Sciences, Huazhong University of Science and Technology, Wuhan 430074, China;

⁹College of Life Sciences, Wuhan University, Wuhan 430072, China;

¹⁰Yunnan Institute of Lamu, Kunming 650034, China;

¹¹School of Science and Information Engineering, Yunnan Agricultural University, Kunming 650201, China;

¹²State Key Laboratory of Genetic Resources and Evolution, Kunming Institute of Zoology, Kunming 650223, China;

¹³Key Laboratory of Pu-er Tea Science, Ministry of Education and Yunnan Agricultural University, Kunming 650201, China

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The drumstick tree (*Moringa oleifera* Lam.) is a perennial crop that has gained popularity in certain developing countries for its high-nutrition content and adaptability to arid and semi-arid environments. Here we report a high-quality draft genome sequence of *M. oleifera*. This assembly represents 91.78% of the estimated genome size and contains 19,465 protein-coding genes. Comparative genomic analysis between *M. oleifera* and related woody plant genomes helps clarify the general evolution of this species, while the identification of several species-specific gene families and positively selected genes in *M. oleifera* may help identify genes related to *M. oleifera*'s high protein content, fast-growth, heat and stress tolerance. This reference genome greatly extends the basic research on *M. oleifera*, and may further promote applying genomics to enhanced breeding and improvement of *M. oleifera*.

genome, drumstick tree, *Moringa oleifera*

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Despite doubling of per-acre yields of major grain crops since 1950, nearly one in seven people suffer from malnu-

trition worldwide, predominately in developing countries. While there are myriad reasons for widespread lack of foodstuffs, part of the problem is that the human food system is dominated by annual crops that are sown each year, such as corn, wheat, rice, and most of leaf vegetables. These

†Contributed equally to this work

*Corresponding author (email: shengj@ynau.edu.cn; haoism@sina.com)

crops can be quite labor and resource intensive, and are not all suited to certain ecosystems found in many developing countries. Conversely, certain perennials—plants that are sown once and live for years—that are highly water-use and nutrient cycling efficient, adaptable to a wide array of environments, have high-nutrition value, may be viable alternatives to traditional annuals. Unfortunately, to date few of these perennial plants are widely planted or consumed, with the banana, coco and pigeon pea being notable exceptions.

Recently, a small to medium-sized, evergreen or deciduous tree native to northern India, Pakistan and Nepal known as the drumstick tree (*Moringa oleifera* Lam.)—or alternatively as the horseradish tree or ben oil tree—has received increased agricultural and industrial attention. Not only can every part of the tree be used as food, medicines or for industrial purposes [1–4], but its high protein, vitamin and mineral content have made it an attractive target for wide-spread planting in some developing countries [1,3,5,6]. Moreover, *M. oleifera* grows well at altitudes from 0 to 1,800 m and in areas with rainfall between 500 and 1,500 mm per year, making it suitable for both semi-arid and arid ecosystem, which covers 37.0% of the earth's geographical area, and even larger swaths of the developing world. Despite these benefits and efforts to cultivate the tree, little basic research on *M. oleifera* has been conducted, which greatly limits its further traditional and novel applications. Here, for the first time we have sequenced the genome of *M. oleifera* and provided well-assembled and annotated genome that should prove invaluable in furthering the uses and investigations of this important perennial.

1 Materials and methods

1.1 DNA materials

DNA used for sequencing was extracted from the leaves of a one-year-old drumstick tree (*Moringa oleifera* Lam.) planted in Pu'er, Yunnan province, China. In total, over 50 μ g DNA was used to construct the sequencing libraries.

1.2 Sequencing data production and processing

Whole genome shotgun sequencing was deployed to produce reads on an Illumina Hiseq2500TM. Raw sequencing data of Illumina Hiseq2500TM were obtained through three steps, image analysis, base calling and sequence analysis, yielded a total of 202 Gb raw data.

Reads with more than 10 low-quality bases (low-quality being defined as value less than 60 bases or with “N”s were filtered). Duplicated reads and reads with adaptor are also removed. Both ends of each read were trimmed by 2 bp. Totally, we obtained seven libraries of different lengths: 177, 222, 390, 503, 3,500, 11,500, and 15,000 bp. Adaptor ligation and DNA cluster preparation were performed prior

to sequencing. *K*-mer frequency and correction to reduce low frequency reads (primarily caused by sequencing errors) was done in SOAPec 2.01 [7], leaving clean data suitable for *de novo* assembly.

1.3 Genome assembly

To deal with genomes with the potential high heterozygosity of the genomes, Platanus 1.2.1 [8] was used to assemble DNA fragments (reads) into contigs. The initial *K*-mer size was set 41, step size was 10, maximum difference for branch cutting was 0.3, maximum difference for bubble crush was 0.15, and *K*-mer coverage cutoff was 5. No parameter needs to be set during scaffolding using SSPACE v2.0 [9]. The final assembly was generated after gap filling with Gapcloser v1.12 in SOAPdenovo package [7]. The assembly was evaluated by mapping the reads back to the genome using SOAPaligner 2.18.

1.4 Repetitive sequence annotation

Tandem Repeats Finder (TRF) 4.04 [10] was used to identify tandem repeats in the *M. oleifera* genome, and then Repeatmasker 3.3.0 and RepeatProteinMask were used to search repeats against Repbase [11] at the DNA and protein levels, respectively. These results were combined with the *de novo* prediction via LTR_FINDER 1.05 [12] and RepeatScout [13].

1.5 Protein-coding gene annotation

A combination of homology-based and *ab initio* methods yielded 19,465 annotated protein-coding genes. Protein sequences of six plant species (*Arabidopsis thaliana* [14], *Glycine max* [15], *Oryza sativa* [16], *Populus trichocarpa* [17], *Sorghum bicolor* [18], *Selaginella moellendorffii* [19]) were used in the homology-based method. These six species were selected because they all have well assembled and annotated genomes sequences, and include angiosperm to gymnosperm species that are related to *M. oleifera*, making them excellent reference points to explore the evolutionary processes related to *M. oleifera*. In the homology-based method, we first performed tblastn setting *e*-value cutoff 10^{-5} . Blast hits with *e*-value lower than 10^{-5} in the genome were discarded, and then predicted regions were extended by 2,000 bp both upstream and downstream, and aligned against protein sequence using GeneWise [20] to identify gene structure. With the AUGUSTUS 2.5.5 [21], Genscan, and GlimmerHMM 3.0.1 [22] software packages used for gene prediction. In the *ab initio* method, the genes predicted by software were aligned to *Arabidopsis thaliana* protein sequences, with alignment rate set at 0.5. The two sets of genes were then merged using GLEAN, a software that can create consensus gene sets by integrating disparate sources of gene structure evidence.

1.6 Gene function annotation

Potential functions of the annotated genes were assigned by choosing the best alignment of genes against the TrEMBL [23], KEGG [24] and InterProscan [25] databases.

1.7 Non-coding gene annotation

tRNAscan-SE v1.23 [26] was used for *M. oleifera* tRNA annotation. We used homology method to identify rRNA. rRNA sequence data was downloaded from the Rfam [27] database to serve as a reference. INFERNAL v0.81 [28] was used to identify snRNA and miRNA.

1.8 miRNA target analysis

Mature miRNA sequences were downloaded from miRbase [29] and aligned to annotated miRNA genes via blastn. Hits longer than 16 bp were selected as potential mature miRNA sequences. Then we predicted target genes of these mature miRNA sequences using online tool psRNATarget [30].

1.9 Gene families

Four other species including *Vitis vinifera*, *Cajanus cajan*, *Carica papaya*, *Malus pumila* and software OrthoMCL 1.4 [31] were used to identify gene clusters. First, we conducted pairwise alignment using blastp with *e*-value cutoff of 10^{-5} . Then OrthoMCL was used with all parameters default.

1.10 Phylogenetic relationship and divergent time

Single copy gene family genes of the five woody plants (*Moringa oleifera*, *Vitis vinifera*, *Cajanus cajan*, *Carica papaya*, *Malus pumila*) obtained from gene family analysis were used for phylogenetic analysis. Multiple sequence alignments were performed using MUSCLE 3.8.31 [32]. Four-fold degenerate sites were extracted from each gene and concatenated into one linear sequence for each species, in order to construct a neighbor joining tree using PhyML 3.0. To estimate the divergence time of each species, we used known divergence time information between plant

species from the public resource, TIMETREE (<http://www.timetree.org/>). Using data generated from the phylogenetic tree, we estimated divergence times with MCMCTREE in paml 4.4 [33].

1.11 Gene family contraction and expanding

CAFE 2.1 [34] was used to screening gene family expansion and contraction history.

1.12 Positive selection analysis

Blast was performed to align the coding sequence data of *M. oleifera* and *Carica papaya* in order to find the gene pairs with the best alignments. The resulting 5,601 orthologous gene pairs were aligned again using lastz as a preparation for KaKs_Calculator 1.2 [35], which finally yielded a dataset of each gene pair's *Ka/Ks* ratio. Alignment in Figure 3 and Figure S10 was produced by multiple alignment tool MUSCLE [32] and picture was generated by ClustalX [36].

2 Results

2.1 Genome assembly of *M. oleifera*

We obtained 457× coverage DNA sequencing data for the *M. oleifera* sample (summary of sequencing data used for the assembly is presented in Table S1, and 17-mer frequency distribution is shown in Figure S1). Based on the 17-mer frequency distribution, the estimated genome size was estimated at 315 Mb (Table S2), and further flow cytometry indicated that the nuclear genome size (*c*-value) of *M. oleifera* was comparable and/or smaller than that of *Oryza sativa*. The final contig and scaffold N50 were 123 kb and 1.14 Mb, respectively (Table 1), with over 80% (231 Mb) of the total sequence represented in 262 scaffolds. The final quality of genome assembly was comparable to recently published high-quality reference plant genomes [19]. In total, 95.67% reads could be re-mapped to the assembly, further confirmed the quality of our genome assembly (Table S3).

The genome size of woody plant ranges from 280 Mb, such as *Prunus mume* [37], to 221.8 Gb for *Pinus taeda* [38].

Table 1 Summary statistics of *M. oleifera* genome assembly

	Contig		Scaffold	
	Size (bp)	Number	Size (bp)	Number
N90	4,165	4,362	5,792	1,382
N80	30,989	1,914	150,929	262
N70	60,562	1,261	396,940	147
N60	91,660	880	736,902	93
N50	123,008	611	1,140,476	61
Longest	1,070,888		6,788,971	
Average size	6,911		8,677	
Total number (>1,000bp)		13512		10,494
Total	287,419,725	41,586		33,332

Here, the genomes of *M. oleifera* proved to be among the smallest, being even smaller than rice. Paired with *M. oleifera*'s fast-growth, high seed production, and adaptation to arid and semi-arid environments, the small size of the *M. oleifera* genome makes it not just an attractive perennial, but a viable model for functional genomic studies aimed at better characterizing the woody plant biology.

2.2 Annotation of *M. oleifera* Genome

A combination of homology and *ab initio* methods allowed us to annotate 19,465 high-confidence protein-coding loci in the *M. oleifera* genome with a mean coding length of 3,354.22 bp and an average of 5.42 exons per gene (Table S4). Further gene structure-based evaluation to confirm the annotation of protein-coding genes (distribution of mRNA structure statistics are in Figures S2 and S3) showed that 93.74% of *M. oleifera* genes have homologs in the TrEMBL protein database, and 72.67% could be classified by Swiss-Prot [23]. In total, 94.01% of the genes have either known homologs or can be functionally classified with InterPro, GO, KEGG, Swiss-Prot or TrEMBL databases [39] (Table S5).

Structure- and homology-based analyses identified 148,820,058 bp repetitive elements, covering most types of plant transposable elements. Most of the repeats were *de novo* predicted. Curiously, only 10.1% of the repeats detected by homologous method, perhaps reflecting phylogenetic distance of *M. oleifera* from other plants with published genomes. Together with numerous truncated repetitive elements, these elements make up 51.45% of the *M. oleifera* genome (Table S6), while 136 Mb of the repeats were transposable elements (TE) that make up 47.10% of the *M. oleifera* genome (Table S7; distribution of TE divergence rate is shown in Figures S4 and S5). An overview of annotated non-coding RNA (ncRNA) genes is shown in Table S8. In total, we predicted 87 mature miRNAs and 369 potential target genes of these miRNAs (Table S9). GO (gene ontology) [40] enrichment analysis of these genes using Ontologizer [41] (Figure S6) showed that 25 of 26 enriched terms were concentrated in cellular biological process regulation.

Previous studies found that intracellular tRNA level may be correlated with tRNA gene copy number [42]. Here, 1,777 tRNA genes reside in *M. oleifera* genome, but only 388 in *Carica papaya* and 600 in *Vitis vinifera*, which may related to *M. oleifera*'s markedly high protein synthesis ability.

2.3 Phylogenetic and whole genome duplication analysis

M. oleifera was originally classified into *Rhoadales* in *Flora of China* based on its morphology, but mounting molecular evidences suggest it belongs to *Brassicales* [43].

Here, four woody plants with published genomes from the *Dicotyledons* clade—*Vitis vinifera* [44], *Cajanus cajan* [45], *Carica papaya* [46], and *Malus domestica* [47]—were used to construct a phylogenetic tree of *M. oleifera* (Figure 1A shows the divergence time of each branch). The tree showed that *Carica papaya* is the most closely related species to *M. oleifera*, suggesting placement of this species in *Brassicales*. And we analyzed the phylogeny of four *Brassicales* (*Arabidopsis thaliana*, *Brassica rapa*, *carica papaya*, and *Moringa oleifera*; shown in Figure S7). Further whole genome duplication analysis of these four *Brassicales* indicated that whole genome duplication (WGD) events took place several times in *Brassicales* (Figure 1C). Such WGD events help clarify some of the history of *Brassicales*; for example, *Carica papaya* was previously known to have not experienced the At- β WGD [46], and our data suggests that neither *M. oleifera* nor *Carica papaya* have experienced any recent WGD events (Figure 1C). Instead, our analyses indicate that the last WGD events of these two species took place before they diverged from *A. thaliana*, that is the At- γ WGD; a finding further supported by calculating the *Ks* between the paralogous genes of *M. oleifera* (Figure 1D) which showed only one obvious peak where *Ks* \approx 1.8.

2.4 *M. oleifera*-specific gene families and genes

Gene family is often an assemblage of genes with approximately the same function. Species-specific gene families add raw materials to the generation of discrepancy against other species [48,49]. Here, we carried out gene family clustering analysis on all protein-coding genes of *M. oleifera*. Comparative analysis of *M. oleifera* with *Vitis vinifera*, *Cajanus cajan*, *Carica papaya* and *Malus domestica* showed that these five different plant species possess similar numbers of gene families, with a core set of 10,215 shared genes (Figure 1b). Compared to other species, however, *M. oleifera* has markedly fewer single-copy families and unclustered genes (distribution of gene clusters in *M. oleifera* genome was shown in Table S10 and Figure S8). Of 12,298 gene families in *M. oleifera*, 198 gene families were *M. oleifera*-specific, including a total of 812 genes (GO enrichment analysis of these genes is in Figure S9). Calculating gene family contraction and expansion on each branch in the phylogenetic tree showed that that *M. oleifera* has 560 expanded gene families—the smallest of the five wood plant species with published genomes—and 2,611 contracted gene families (Figure 1A), making it comparatively smaller with a compact genome.

Curiously, four *SKP1* genes and 18 F-box domain containing genes were identified as members of *M. oleifera*-specific families. Gene *SKP1* is a protein crucial to the cell cycle controlling [50] that helps coordinated the ubiquitination and degradation of phase specific proteins to maintain the cell cycle, with the F-box motif maintaining the association between these proteins [51]. Rather confusingly,

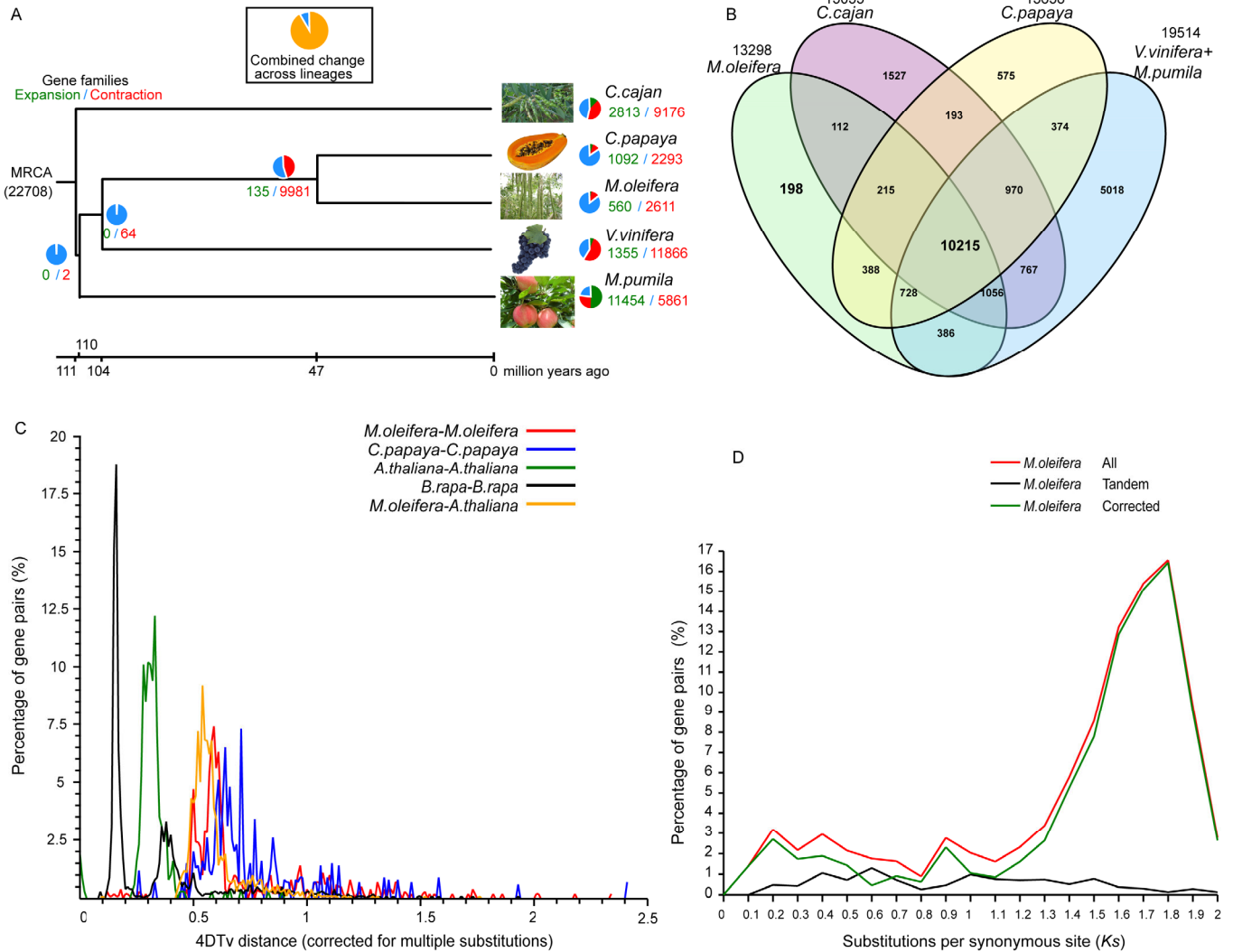


Figure 1 Comparative genomic analysis. A, Phylogenetic tree of five woody plants, *M. oleifera*, *Vitis vinifera*, *Cajanus cajan*, *Carica papaya* and *Malus domestica*. Estimated divergence time ranges from 47 million years ago (mya) to 110 mya. Numbers of gene family contractions and expansions are shown in the pie chart. B, Gene family analysis of *M. oleifera* against *Vitis vinifera*, *Cajanus cajan*, *Carica papaya* and *Malus domestica*. *Vitis vinifera* and *Malus domestica* were merged into one dataset. C, 4Dtv (fourfold degenerate third-codon transversion) analysis of *M. oleifera* using *Arabidopsis thaliana*, *Brassica rapa* and *Carica papaya*. D, Distribution of *Ks* between paralogous gene pairs.

four *SKP1* and 18 F-box containing genes *M. oleifera* were in the *M. oleifera*-specific gene families while another seven *SKP1* genes and 104 F-box containing genes were not. In theory, there may be two potential explanations: First, these represented genes may be newly derived and may play a role in *M. oleifera*'s fast-growth and heat tolerance, while second these genes may simply be redundant and accumulated many mutations. We also found three *BET V 1* genes in the *M. oleifera*-specific gene families. *BET V 1* was first found in birch tree pollen as an allergen [52], but more functions of this gene have been discovered later, including its role as a steroid carrier [53]. *BET V 1* genes are potential factors for the *M. oleifera*'s fast growth as this gene family is related to the binding of many ligands, including ABA, lipids and steroids. These *M. oleifera*-specific genes may be functionally important to *M. oleifera* and warrant further

investigation.

2.5 Positively selected genes in *M. oleifera* genome

Positively selected genes often have functions that favor the organism's adaptation and establishment in an area. To investigate which genes may be associated with certain traits that have made *M. oleifera*'s successful, we conducted positive selection analysis. Blast and KaKs_Calculator [54] compared orthologs between *M. oleifera* and each one of *Carica papaya*, *Vitis vinifera* and *Malus domestica*, and respectively found 566, 399, 112 genes of *M. oleifera* with *Ka/Ks* ratio > 1 (significance, $P < 0.05$; see Table S11, Table S12, Table S13). We further found four genes that overlap among the three gene sets (Figure 2). We also found two genes (annotated gene: *lamu_GLEAN_10016878*, *lamu_*

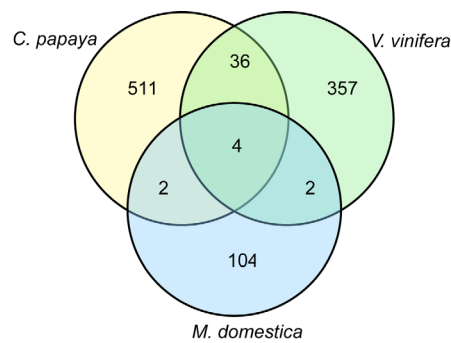


Figure 2 Positively selected genes identified from pairwise comparisons between *M. oleifera* and *Carica papaya*, *Vitis vinifera* and *Malus domestica*.

GLEAN_10011614) with alignment length longer than half the gene size, indicating they may be specifically and strongly selected in *M. oleifera*. Gene *lamu_GLEAN_10016878* is functionally annotated as a Myb/SANT-like DNA binding domains. The SANT domain is ubiquitous in chromatin regulatory proteins and it is often involved in histone acetylation, deacetylation and ATP-dependent chromatin remodeling process [55]. More importantly, many proteins containing the Myb/SANT domain have DNA binding activity and are related to gene regulation; these regulators of different functions often do not resemble each other. However, both ends of this gene and some segments of the central region is highly conserved. However some parts of central regions vary (see screening of the alignment of orthologous genes in Figure S10) [56,57].

Gene *lamu_GLEAN_10011614* is supposed to be a ribosomal protein S6e, which is highly conserved in vertebrates, invertebrates, and fungi [58]. In eukaryotes, ribosomal proteins synthesized in cytoplasm are imported into nucleoplasm before associating with newly transcribed pre-rRNA to form a 90S complex, which is processed into a 60S and a 40S ribosomal subunit and subsequently exported into cyto-

plasm [59]. Ribosomal proteins assist the maturation and functioning of pre-18S RNA and ribosome [60]. The S6e amino acid sequence has twonucleolar binding sequence (Nobis) and severalnuclear localization signal (NLS) sequence according to Kundu-Michalik's study [58]. We identified Nobis1's N-terminus by recognizing (G)RVRL pattern and inferred the C-terminus according to the length of Nobis1 revealed in previous study [58]. Based on the Kundu-Michalik's study, we also proposed a rough border of the Nobis2 frame. And some other elements such as NLS and phosphorylation sites on this sequence were tentatively presented in Figure 3. Phosphorylation states of S6 often act as a switch and regulate cell processes through phosphorylation cascade [61]. Positive selection of the *lamu_GLEAN_10011614* gene elegantly serves as further molecular evidence that the protein synthesis machinery of *M. oleifera* has likely experienced strong evolutionary rewiring to produce more proteins.

2.6 Analysis of transcription factor families

Transcription factors regulate gene expression, making them crucial and diverse in organisms ranging from microbes to high plants and animals [62,63]. Analysis of previously identified transcription factors has yielded a large amount of information on gene expression patterns. Taking the *Arabidopsis thaliana* transcription factor families in the TAIR database (<http://arabidopsis.org/browse/genefamily/index.jsp>) [64] as reference, we identified a total of 939 transcription factors (Table S14) in the *M. oleifera* genome using the blastp with $P\text{-value} < 10^{-20}$. Interestingly, our positive selection analysis of *M. oleifera* against each of *Vitis vinifera*, *Carica papaya*, and *Malus domestica* uncovered 43 transcription factors were under positive selection in various classes including ABI3VP1, AP2-EREBP, Alfin-like, C2C2-Dof, C2C2-Gata, C2H2, C3H, CPP, E2F-DP,

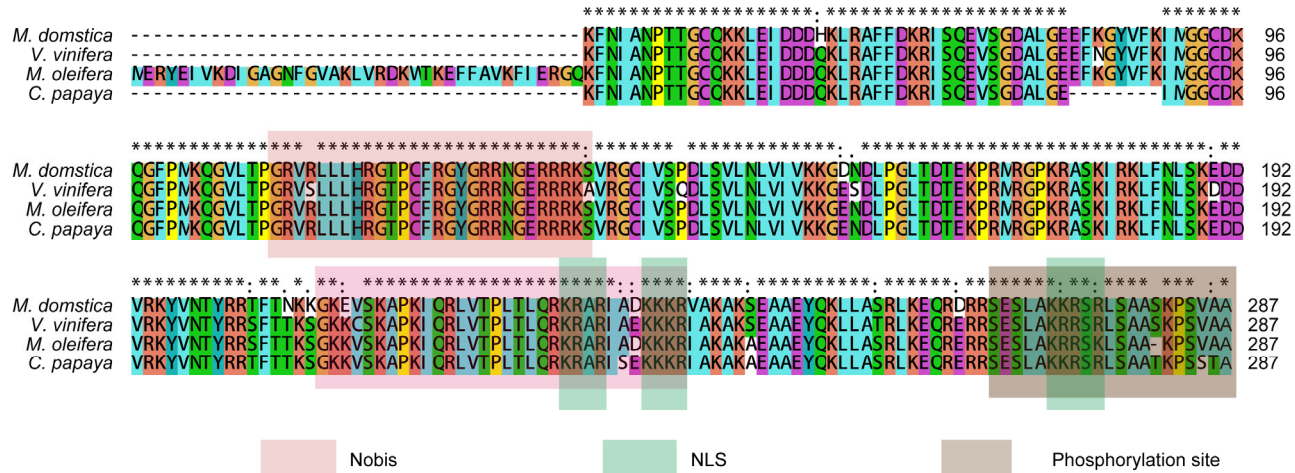


Figure 3 Alignment of *M. oleifera* gene *lamu_GLEAN_10011614* and its orthologous in *Vitis vinifera*, *Cajanus cajan*, *Carica papaya*, *Malus domestica*. Postulated elements like Nobis, NLS, and phosphorylation sites have been marked in colored rectangles.

G2-like, GRAS, Homeobox, MADS, MYB, NAC, PHD, Trihelix, WRKY, bHLH transcription factors. Among these, the WRKY transcription factors are particularly interesting, since they were previously suggested to play important roles in response to various abiotic stress, including cold, heat, water deficiency, excessive salt, nutrient starvation, and variable light condition. Here, we found five copies under positive selection. Similarly, the C2H2 transcription factors—a superfamily that plays important roles in defense responses and various other physiological processes in plants—have four copies under positive selection. Meanwhile, the AP2-EREBP transcription factors that were previously implicated in hormone, sugar and redox signaling in context of abiotic stresses such as cold and drought, had two copies under positive selection, while the C3H transcription factors with some copies being reported to response to drought stress have two copies under positive selection (Table S15). The fact that all of these transcription factors involved in stress response were found to be under positive selection may account for *M. oleifera*'s adaptation to both heat and drought stress present in arid environments.

2.7 HSP genes

Heat stress is a serious threat to crop production, which may be exacerbated by changes in global climates. Accordingly, the high temperature tolerance of *M. oleifera* [1] may prove quite useful. In many species of plants, Heat shock proteins (HSPs) or stress-induced proteins participate in many primary stress response such as drought, salinity, cold and hot temperatures and chemicals [65–67]. Using the heat shock proteins sequences of *Arabidopsis thaliana* we downloaded from HSPiR (<http://pds-lab.biochem.iisc.ernet.in/hspir/chaperone.php>) [68] as reference, we identified a total of 133 heat shock proteins. Based on their nature of functions and molecular mass, HSPs are classified broadly into six major families, namely Hsp70 (25 copies in *M. oleifera* genome), Hsp40 (J-proteins, 52 copies in *M. oleifera* genome), Hsp60 (chaperonins, 17 copies in *M. oleifera* genome), Hsp90 (three copies in *M. oleifera* genome), Hsp100 (Clp proteins, nine copies in *M. oleifera* genome) and small heat shock proteins (27 sHsps copies in *M. oleifera* genome) (Table S16).

We further checked the HSP genes' *Ka/Ks* ratio between *M. oleifera* and *Carica papaya*, and found that the average *Ka/Ks* ratio of HSP genes was higher than that of the background (Table S17). HSP genes that have positive selection features against any of *Carica papaya*, *Vitis vinifera* and *Malus domestica* were collected and shown in Table S18. These genes may potentially be related to the heat tolerance that is one characteristic of *M. oleifera*.

2.8 Brassinosteroid signal transduction pathway

Brassinosteroid is a kind of plant hormone with a regulatory

function in cell elongation and cell division, which can significantly promote plant growth. Previous reports suggest that brassinosteroids help plants get through environmental stresses such as cold, drought and heat. Here, we analyzed the brassinosteroid signal transduction pathway in *M. oleifera* and found that the *BAK1* (BRI1 associated receptor kinase 1) gene expanded in *M. oleifera* with 29 copies, as compared to five copies in *A. thaliana* genome (Figure S11). Furthermore, we also noticed one copy of the *BAK1* gene was also under positive selection when compared against *Vitis vinifera*. *BAK1* plays a major role in transducing the BR signal, and loss-of-function mutation of *BAK1* caused a weak dwarf phenotype [69].

2.9 γ -aminobutyrate (GABA) bio-synthesis and sitosterol bio-synthesis pathways in *M. oleifera*

We analyzed GABA bio-synthesis and sitosterol bio-synthesis pathways in *M. oleifera*—both of which are important hormone pathways in plants—and annotated all genes in the pathways. 4-Aminobutyrate or GABA is a ubiquitous, four carbon, non-protein amino acid found in higher plants, animals, fungi and bacteria. In plants, the concentration of GABA is markedly stimulated by a variety of stress conditions, *e.g.* hypoxia, temperature shock, mechanical manipulation and damage, water stress and phytohormones [70,71]. Here, we found that GABA is synthesized almost exclusively by the irreversible α -decarboxylation of L-glutamate by glutamate decarboxylase (GAD; annotated gene: *lamu_GLEAN_10006873*, *lamu_GLEAN_10006874*, *lamu_GLEAN_10004957*, *lamu_GLEAN_10007711*, *lamu_GLEAN_10007712*, *lamu_GLEAN_10007713*) [72,73]. Subsequently, GABA is catabolized by GABA transaminase (GABA-T; annotated gene: *lamu_GLEAN_10002543*) and succinate semialdehyde dehydrogenase (SSADH; annotated gene: *lamu_GLEAN_10008793*, *lamu_GLEAN_10008794*) to succinate, an important Krebs cycle metabolite [73]. The only other enzyme of glutamate metabolism known to be stimulated by Ca^{2+} in plants is glutamate dehydrogenase (GDH; annotated gene: *lamu_GLEAN_10005665*), a mitochondrial enzyme (Figure S12).

To understand the sterol biosynthesis genes in *M. oleifera*, we tried to draw the major sterol biosynthetic pathway operating in most higher plants [74]. Sitosterol is a typical plant membrane reinforcement, at the expense of campesterol [75]. Campesterol can be used to produce brassinosteroids, which were reported to have observable growth-promoting effects in many plants. The gene *STM2*, of which two copies were found in *M. oleifera*, plays a critical role in balancing the ratio of campesterol to sitosterol to satisfy both growth requirements and membrane integrity (Figure S13).

3 Discussion

At the time of this study, no genomes of species in the family *Moringaceae* were available, making the present *M. oleifera* genome data a valuable reference for further studies on both *M. oleifera* and other species in this important plant family. Due to a dearth of related research, this present study is far from conclusive on many fronts, and is instead suggestive of many further lines of inquiry into the unique characteristics of *M. oleifera* that remain to be explored. In particular, the gene cluster analysis reveals that *M. oleifera* possesses a remarkably small amount of single copy genes, and small amount of *M. oleifera* specific gene families. Taken alongside the fact that the annotated *M. oleifera* genes were fewer than any other resolved higher plants indicate that *M. oleifera* has a compact genome, which may, in part, be responsible or underlie its comparatively fast growth and rapid cell proliferation.

In the present study, we concentrated on the indistinct relationship between the genome content characters and the phenotypic traits, identified a number of genes or gene families that might account for the high protein content, heat tolerance, drought resistance, and fast growth of *M. oleifera*. The gene list provided by our analysis is important not only for the future functional studies of *M. oleifera*, but also for future efforts in breeding and improvement of *M. oleifera*, both of which may help promote *M. oleifera* as a viable perennial crop in regions of the world where food shortages are endemic or the local environment cannot support more traditional annual crops.

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Supporting Information

Figure S1 17-mer frequency distribution of reads data. A minor peak is at about half the depth of the main peak, indicating the heterozygosity of *M. oleifera*.

Figure S2 Distribution of exon numbers in annotated mRNA sequences. Annotation data of *Vitis vinifera*, *Populus trichocarpa*, *Selaginella moellendorffii* were used in parallel with *Moringa oleifera*'s annotation data.

Figure S3 Distribution of mRNA length, CDS length, exon length and intron length. Alongside the genome annotation of *Moringa oleifera*, we used *Vitis vinifera*, *Populus trichocarpa*, *Selaginella moellendorffii* genome annotation as references.

Figure S4 Distribution of Divergence Rate of each Type of *Moringa oleifera*'s TE. Divergence rate was calculated between the identified TE elements in the genome by homology-based method and the consensus sequence in the Repbase.

Figure S5 Distribution of Divergence Rate of each Type of *Moringa oleifera*'s TE. Divergence rate was calculated between the identified TE elements in the genome by de novo method and the consensus sequence in the predicted TE library.

Figure S6 GO analysis of the miRNA targeted gene. Note: *P*-value cutoff was set at 0.0005 to optimize the image.

Figure S7 Phylogenetic tree of four Brassicales (*Arabidopsis thaliana*, *Brassica rapa*, *Carica papaya*, and *Moringa oleifera*).

Figure S8 Orthologous gene distribution among five woody plants. *Vitis vinifera*, *Cajanus cajan*, *Carica papaya*, *Malus pumila* serve as the out group species to search the orthologous genes.

Figure S9 GO enrichment analysis of the *M. oleifera* specific gene family. These genes belong to the family that only exists in *M. oleifera* but not in *Vitis vinifera*, *Cajanus cajan*, *Carica papaya*, *Malus pumila*. *P*-value cutoff is set 0.01.

Figure S10 Alignment of *M. oleifera* gene *lamu_GLEAN_10011614* and its orthologous in *Vitis vinifera*, *Cajanus cajan*, *Carica papaya*, *Malus domestica*. The central region has more variations and N terminal is highly conserved.

Figure S11 Brassinosteroid signal transduction pathway in *M. oleifera* and *Arabidopsis thaliana*. The number near the left square brackets indicates the copies of this gene in the *Arabidopsis thaliana* genome, while the other indicates the copy number in the *M. oleifera* genome.

Figure S12 Simplified metabolic diagram the GABA shunt in relation to the Krebs cycle. GAD, glutamate decarboxylase; GABA-T, GABA transaminase; GDH, glutamate dehydrogenase; SSADH, succinic semialdehyde dehydrogenase.

Figure S13 Biosynthesis of (24 ζ)-24-methyl cholesterol (campesterol) and (24R)-24-ethyl cholesterol (sitosterol) in *Arabidopsis thaliana* and *M. oleifera* genome. CPI, cyclopropyl sterol isomerase; SMT, sterol methyltransferase; OBT14DM, obtusifoliol-14-demethylase; SMO, sterol 4-methyl oxidase; DWF1, gene encoding the Δ 5-sterol- Δ 24-reductase. (isomerase); FACKEL, gene encoding the Δ 8,14-sterol- Δ 14-reductase; HYDRA1, gene encoding the Δ 8- Δ 7-sterol isomerase; DWF5, gene encoding the Δ 5,7-sterol- Δ 7-reductase; DWF7, Δ 7-sterol-C5(6)-desaturase.

Table S1 Statistics of raw data

Table S2 Statistics of 17-mer analysis

Table S3 Statistics of mapping reads to the genome assembly

Table S4 Statistics of gene annotation

Table S5 Overview of gene function annotation.

Table S6 Statistics of Repeats in *M. oleifera* Genome

Table S7 TEs Content in the Assembled *Moringa oleifera* Genome

Table S8 Annotated ncRNA in the genome

Table S9 Predicted miRNA target genes

Table S10 Overview of gene family clustering among *M. oleifera* and *V. vinifera*, *C. cajan*, *C. papaya*, *M. pumila*

Table S11 Positively selected genes of *M. oleifera* against *Carica papaya*

Table S12 Positively selected genes of *M. oleifera* against *Vitis vinifera*

Table S13 Positively selected genes of *M. oleifera* against *Malus pumila*

Table S14 Identified transcription factors

Table S15 Positively selected transcription factor genes in *M. oleifera*

Table S16 Heat Shock Proteins (HSPs) in the *M. oleifera* genome

Table S17 HSP genes' *Ka/Ks* ratio and a comparison with the background

Table S18 Positively selected HSP genes in *M. oleifera*

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